



```
FT Modified-site /note= "potential phosphorylation site"
FT 284
FT /note= "potential phosphorylation site"
FT 285
FT Modified-site /note= "potential phosphorylation site"
FT 295
FT /note= "potential phosphorylation site"
FT
XX
XX WO200023589-A2.
XX
XX
XX 27-APR-2000.
XX
XX 19-OCT-1999; 99WO-US24511.
XX
XX PF 20-OCT-1998; 98US-0172216.
XX
XX PR 04-FEB-1999; 99US-0118559.
XX
XX PR 11-FEB-1999; 99US-0172229.
XX
XX PR 22-APR-1999; 99US-0154336.
XX
XX (INCY-) INCYTE PHARM INC.
XX
XX Tang YT, Yue H, Hillman JL, Guegler KJ, Corley NC, Lal P;
XX PI Azlmzal Y, Baughn MR, Yang J, Shih LL.
XX
XX WPI: 2000-339688/29.
XX
XX N-PSDB: AAA14991.
XX
XX
XX New human proliferation and apoptosis related protein polypeptides used
XX PT for diagnosis, treatment and prevention of cell proliferative,
XX PT immunological and reproductive disorders -
XX
XX Claim 1; Page 89-90; 128pp; English.
XX
XX
XX The present sequence represents a human proliferation and apoptosis
XX CC related protein (PROAP). The polypeptides and polynucleotides can be
XX CC used for the diagnosis, treatment and prevention of cell proliferative,
XX CC immunological and reproductive disorders. Disorders associated with
XX CC decreased expression or activity of include arteriosclerosis, cirrhosis,
XX CC hepatitis, psoriasis, melanoma, lymphoma and cancers of the breast,
XX CC brain and prostate, acquired immune deficiency syndrome (AIDS),
XX CC allergies, anemias, asthma, diabetes mellitus, osteoarthritis,
XX CC endometriosis, uterine fibroids and disruptions of the menstrual cycle.
XX CC Antibodies against PROAP can be used in diagnosis of disorders
XX CC characterized by PROAP e.g. in ELISA (enzyme linked immunosorbent
XX CC assays) and the polynucleotides may be used to detect and quantify gene
XX CC expression in biopsied tissues. These techniques can also be used to
XX CC monitor regulation of PROAP levels during therapeutic intervention.
XX
XX
XX Sequence 281 AA:
SQ
Query Match 100.0%; Score 1492; DB 21; Length 281;
Best Local Similarity 100.0%; Pred. No. 2,76-146;
Matches 281; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
```

```
RESULT 2
AAW01750
ID AAW01750 standard; Protein; 268 AA.
XX
XX AAW01750;
XX
XX 04-SEP-1997 (first entry)
XX
XX
XX ECORI binding fragment protein 1 (EB1).
XX
XX
XX EBI, ECORI Binding Fragment protein 1; cellular; APC; truncation;
XX KM adenomatous polyposis coli; tumour suppressor; sporadic; familial;
XX KM colorectal cancer; predisposition; diagnosis; neoplasm; treatment.
XX
XX Homo sapiens.
XX
XX CWO9637611-AL.
XX
XX 28-NOV-1996.
XX
XX 22-MAY-1996; 96WO-US07747.
XX
XX 22-MAY-1995; 95US-0446919.
XX
XX (UYXO ) UNITV JOHNS HOPKINS.
XX
XX Kinzler K, Vogelstein B, Kinzler K;
XX
XX WPI: 1997-021220/02.
XX
XX N-PSDB: AAT59331.
XX
XX
XX EBI DNA and polypeptide(s) - used to determine a pre-disposition to
XX PT or diagnose neoplasms and to assess treatment options.
XX
XX Claim 5; Page 19-21; 45pp; English.
XX
XX
XX This sequence is that of an EBI protein (ECORI Binding Fragment protein
XX CC 1). This cellular protein associates with the carboxyl terminus of APC
XX CC (adenomatous polyposis coli) tumour suppressor gene product. The APC
XX CC tumour suppressor gene plays an important role in the development of both
XX CC sporadic and familial forms of colorectal cancers. Because most APC
XX CC mutations result in the truncation of the APC protein, these mutant APC
XX CC proteins cannot associate with EBI. This suggests that the interaction
XX CC between APC and EBI is important for the normal function of APC and that
XX CC loss of this association is essential for the development of colorectal
XX CC cancer. By assaying for the presence of APC-EBI protein complexes in a
XX CC cell, a predisposition to or diagnosis of neoplasms can be determined.
XX CC EBI can also be used to assess treatment options for cancer cells (which
XX CC are good candidates for treatment with cycloooxygenase inhibitors).
XX
XX
XX Sequence 268 AA:
SQ
Query Match 61.0%; Score 909.5; DB 18; Length 268;
Best Local Similarity 64.6%; Pred. No. 86-86;
Matches 186; Conservative 28; Mismatches 47; Indels 27; Gaps 6;
```

QY 234 ELICQEHSENSPVISGILYATEEGFAPPEDEIEEHOEDDEX 281  
 |||||:| |||:| :| |||||:| | :| |||:| |  
 Db 225 ELICQENGENDPVLQRIYDILYATDEGFVLPD----EGFQEEDEEY 268

RESULT 3  
 ABB57105  
 ID ABB57105 standard; Protein: 268 AA.

AC ABB57105;

DT 07-MAR-2002 (first entry)

DE Mouse ischaemic condition related protein sequence SEQ ID NO:239.

KW Mouse; ischaemia; compressive ischaemia; occlusive ischaemia;  
 KW vasospastic ischaemia; ischaemic condition; ischaemic disease.

OS Mus musculus.

PN W0200188188-A2.

PD 22-NOV-2001.

PE 18-MAY-2001; 2001WO-JP04192.

PR 18-MAY-2000; 2000JP-0145977.

PA (UYNI-) UNIV NIHON SCHOOL JURIDICAL PERSON.

PI Ishikawa K, Asai S, Takahashi Y, Nagata T, Ishii Y;

DR WPI: 2002-034733/04.

DR N-PSDB: AB199334.

PT Examining the ischemic condition (e.g. occlusive ischemia) by measuring  
 PT expression levels of particular genes defined in the specification or  
 PT by determining the expression profile of a gene group comprising these  
 PT genes -

PS Claim 2; Page 676-677; 2690pp; English.

XX The present invention describes a method for examining ischaemic  
 CC conditions, comprising measuring the expression levels of particular  
 CC genes (I) in a test sample or determining the expression profile of a  
 CC gene group in the sample comprising genes selected from (I). The method  
 CC is useful for examining the ischaemic condition (e.g. compressive  
 CC ischaemia, occlusive ischaemia or vasospastic ischaemia) by measuring  
 CC expression levels of particular genes (AB199202 to AB199912, encoding  
 CC the protein sequences in ABB57020 to ABB57374) or by determining the  
 CC expression profile of a gene group comprising these genes. The  
 CC expression levels or expression profiles produced by these genes are  
 CC used as an indicator when screening for ischaemic condition-improving  
 CC drugs or therapeutics for ischaemic diseases. AB199913 and AB199914  
 CC represent PCR primers for a mouse ischaemic condition related sequence,  
 CC which are used in the exemplification of the present invention.

XX Sequence 268 AA;

Query Match 61.0%; Score 909.5; DB 23; Length 268;  
 Best Local Similarity 64.6%; Pred. No. 8e-86;  
 Matches 186; Conservative 29; Mismatches 46; Indels 27; Gaps 6;

QY 1 MAVVNYSTSVSENLSDMLAMVDSLHLNLTKEQLCSGANYQCFMDMLFPGCVHLRK 60  
 |||||:| |||:| :| |||||:| | :| |||:| |

Db 1 MAVVNYSTSVSDNLSDMLAMINESQLNLTKIEQLCSGANYQCFMDMLFPGSIALKK 60  
 |||||:| |||:| :| |||||:| | :| |||:| |

QY VKFOAKLHEHYHNFVLOAAFKKGVCKIIPVEKLVGKPDNDNEFLQFMEKKFEDAND 120  
 |||||:| |||:| :| |||||:| | :| |||:| |

Db 61 VKFOAKLHEHYHNFVLOAAFKKGVCKIIPVEKLVGKPDNDNEFLQFMEKKFEDAND 120  
 |||||:| |||:| :| |||||:| | :| |||:| |

QY 121 GKDYNPLLAROGODVAPPNDQIFNKSKLLIG--TAVPORTSPT-----GPKNMQTS 173

Db 121 GREYDPAVARQOQETFAVAAPSLVAPALSKPKKPLGSSFAAPQAPIMQRTTAPK----- 174  
 |||||:| |||:| :| |||||:| | :| |||:| |

QY 174 RLSNVAPPCILIRKNPPSARANGSHETDAQILELNOQLVDLKLTVDLGKERDPEFSKLRDI 233  
 |||||:| |||:| :| |||||:| | :| |||:| |

Db 175 -----AGPGMWAKN--PGVNG----DDEAAELMQQVKKVLKTVDELKERDPEFSKLRNI 224  
 |||||:| |||:| :| |||||:| | :| |||:| |

QY 234 ELICQEHSENSPVISGILYATEEGFAPPEDEIEEHOEDDEX 281  
 |||||:| |||:| :| |||||:| | :| |||:| |

Db 225 ELICQENGENDPVLQRIYDILYATDEGFVLPD----EGFQEEDEEY 268

RESULT 4  
 AAB43793  
 ID AAB43793 standard; Protein: 311 AA.

AC AAB43793;

DT 08-FEB-2001 (first entry)

DE Human cancer associated protein sequence SEQ ID NO:1238.

XX Homo sapiens.

PN W0200055350-A1.

PD 21-SEP-2000.

PE 08-MAR-2000; 2000WO-US05882.

PR 12-MAR-1999; 99US-0124270.

PA (HUMA-) HUMAN GENOME SCI INC.

PI Rosen CA, Ruben SM;

DR WPI: 2000-587533/55.

DR N-PSDB: AAC78002.

PT Novel isolated nucleic acids comprising sequences encoding peptides  
 PT useful for treating or diagnosing e.g. cancer -

PS Claim 11; Page 1868-1869; 2352pp; English.

XX AAC78002 to AAC78448 encode the human cancer associated proteins given  
 CC in AAB43398 to AAB44239. The proteins can have activities based on the  
 CC tissues and cells the genes are expressed in. Example of activities  
 CC include: cytostatic; proliferative; vulnerable; immunomodulator;  
 CC antiadipetic; antiasthmatic; antirheumatic; antiarthritic;  
 CC antiinflammatory; antihypertensive; antidiabetic; antibacterial;  
 CC antineoplastic; neuroprotective; cardiact; thrombolytic; coagulant;  
 CC neotrophic; vasotropic; antipsoriatic and antiangiogenic. The  
 CC polynucleotides and polypeptides can be used for preventing, treating or  
 CC ameliorating medical conditions and diagnosing pathological conditions.  
 CC Polynucleotides, polypeptides, antibodies, agonists and antagonists from  
 CC the present invention may be used to treat immune disorders by activating  
 CC or inhibiting the proliferation, differentiation or mobilisation of  
 CC immune cells, to treat disorders of haematopoietic cells, autoimmune  
 CC disorders, allergic reactions, graft versus host disease and organ  
 CC rejection, modulate haemostatic or thrombolytic activity, modulate  
 CC inflammation, cancers, cardiovascular disorders, neurological disease and  
 CC bacterial or viral infections. The peptides, nucleotides, antibodies,

CC agonists and antagonists may be also be used in drug screens. AAC78449 to  
CC AAC78457 and AAB44240 represent sequences used in the exemplification of  
CC the present invention.

XX Sequence 311 AA:

Query Match 61.0%; Score 909.5; DB 21; Length 311;  
Best Local Similarity 64.6%; Pred. No. 9,9e-86;  
Matches 186; Conservative 28; Mismatches 47; Indels 27; Gaps 6;

QY 1 MAVNVSTSVTSNTLSRHMLAVNDLHNTYKIEQLCSGAAYCOFMDLPGCVHLRK 60  
DB 44 MAVNVSTSVTSNTLSRHMLAVNDLHNTYKIEQLCSGAAYCOFMDLPGCVHLRK 103  
QY 61 VKFOAKLEHEYIHNKRVLOAARFKMGVDKIIPYEKLVKGFQDNFEFIQWFKKFPDAND 120  
DB 104 VKFOAKLEHEYIHNKRVLOAARFKMGVDKIIPYEKLVKGFQDNFEFIQWFKKFPDAND 163  
QY 121 GADYNPLAROGODVAPPPNPGDQIFNKSCLKI--GTAVPQRTSPT-----GPKNNQTSG 173  
DB 164 GADYDPAAROGODVAPPPNPGDQIFNKSCLKI--GTAVPQRTSPT-----GPKNNQTSG 217  
QY 174 RLSNVAPCILRKNPSPARRNGHETDAQILELNOQLVDLKLTVDGLEKERDFYFSKLRI 233  
DB 218 -----AGPGVVKRN-PGVNG-----DDEAELMQVNVKLTVDELEKERDFYFSKLRI 267  
QY 234 ELICQHESENSPVISGIILYATEEGFAPPEDEIEHQEDODEY 281  
DB 268 ELICQHESENSPVISGIILYATEEGFAPPEDEIEHQEDODEY 311

## RESULT 5

AAG73870  
ID AAG73870 standard; Protein; 311 AA.

XX AAG73870;

XX 03-SEP-2001 (first entry)

DE Human colon cancer antigen protein SEQ ID NO:4634.

KM Human; colon cancer; colon cancer antigen; diagnosis; detection;  
KM colorectal carcinoma; chromosome 20.

XX Homo sapiens.

PN W0200122920-A2.

XX 05-APR-2001.

PF 28-SEP-2000; 2000MO-US26524.

PR 29-SEP-1999; 99US-0157137.

PR 03-NOV-1999; 99US-0163280.

XX (HUMA-) HUMAN GENOME SCI INC.

XX Ruben SM, Barash SC, Blise CF, Rosen CA;

XX MPI. 2001-235357/24.

DR N-PSDB; AAH33301.

XX Nucleic acids encoding 4277 human colon cancer-associated polypeptides,  
XX useful for preventing, diagnosing and/or treating colorectal cancers -  
PS Claim 11; Page 6436-6437; 9803pp; English.

XX AAH32943 to AAH37195 and AAG73514 to AAG77788 represent human colon  
CC cancer-associated nucleic acid molecules (N) and proteins (P), where  
CC the proteins are collectively known as colon cancer antigens. The colon  
CC cancer antigens have cytostatic activity and can be used in gene  
CC therapy and vaccine production. N and P may be used in the prevention,  
CC diagnosis and treatment of diseases associated with inappropriate P

CC expression. For example, N and P may be used to treat disorders  
CC associated with decreased expression by rectifying mutations or deletions  
CC in a patient's genome that affect the activity of P by expressing  
CC inactive proteins or to supplement the patient's own production of P.  
CC Additionally, N may be used to produce the colon cancer-associated Ps,  
CC by inserting the nucleic acids into a host cell and culturing the cell  
CC to express the proteins. N and P can be used in the prevention, diagnosis  
CC and treatment of colorectal carcinomas and cancers. AAH37196 to AAH37204  
CC present invention.

CC N.B. Pages 666 to 682 and page 7053 of the sequence listing were  
CC missing at time of publication, meaning no sequences are present for  
CC SEQ ID NO:1027 to 1052, 7921 and 7922.

XX Sequence 311 AA:

Query Match 61.0%; Score 909.5; DB 22; Length 311;  
Best Local Similarity 64.6%; Pred. No. 9,9e-86;  
Matches 186; Conservative 28; Mismatches 47; Indels 27; Gaps 6;

QY 1 MAVNVSTSVTSNTLSRHMLAVNDLHNTYKIEQLCSGAAYCOFMDLPGCVHLRK 60  
DB 44 MAVNVSTSVTSNTLSRHMLAVNDLHNTYKIEQLCSGAAYCOFMDLPGCVHLRK 103  
QY 61 VKFOAKLEHEYIHNKRVLOAARFKMGVDKIIPYEKLVKGFQDNFEFIQWFKKFPDAND 120  
DB 104 VKFOAKLEHEYIHNKRVLOAARFKMGVDKIIPYEKLVKGFQDNFEFIQWFKKFPDAND 163  
QY 121 GADYNPLAROGODVAPPPNPGDQIFNKSCLKI--GTAVPQRTSPT-----GPKNNQTSG 173  
DB 164 GADYDPAAROGODVAPPPNPGDQIFNKSCLKI--GTAVPQRTSPT-----GPKNNQTSG 217  
QY 174 RLSNVAPCILRKNPSPARRNGHETDAQILELNOQLVDLKLTVDGLEKERDFYFSKLRI 233  
DB 218 -----AGPGVVKRN-PGVNG-----DDEAELMQVNVKLTVDELEKERDFYFSKLRI 267  
QY 234 ELICQHESENSPVISGIILYATEEGFAPPEDEIEHQEDODEY 281  
DB 268 ELICQHESENSPVISGIILYATEEGFAPPEDEIEHQEDODEY 311

## RESULT 6

AAW48626  
ID AAW48626 standard; Protein; 327 AA.

XX AAW48626;

XX 17-AUG-1998 (first entry)

DE Human adenomatous polyposis coli protein-binding protein RPL.

KM RPL; human; adenomatous polyposis coli; APC; T cell activation;

KM lymphoma; colorectal cancer; ulcerative colitis; Crohn's disease;

XX diagnosis.

XX Homo sapiens.

XX Key

XX Location/Qualifiers

FT Misc-difference 60 /note= "encoded by ATC"

FT Misc-difference 61 /note= "encoded by ATT"

FT Misc-difference 62 /note= "encoded by GCA"

FT Misc-difference 63 /note= "encoded by TGG"

FT Misc-difference 64 /note= "encoded by GTT"

FT Misc-difference 65 /note= "encoded by AAT"

FT Misc-difference 66 /note= "encoded by GAC"

FT Misc-difference 67

FT /note= "encoded by ATA"  
 FT Misc-difference 68  
 FT /note= "encoded by GTA"  
 FT Misc-difference 69  
 FT /note= "encoded by TCT"  
 FT Misc-difference 70  
 FT /note= "encoded by AGA"  
 FT Misc-difference 71  
 FT /note= "encoded by CAT"  
 FT Misc-difference 72  
 FT /note= "encoded by GAC"  
 XX  
 PN WO9807737-A1.  
 XX  
 PD 26-FEB-1998.  
 XX  
 PF 21-AUG-1997; 97WO-US14753.  
 XX  
 PR 21-AUG-1996; 96US-0701233.  
 XX  
 PA (LUDW-) LUDWIG INST CANCER RES.  
 XX  
 PI Pfeundschnub M, Renner C;  
 XX  
 DR WPI: 1998-169085/15.  
 DR N-PSDB: AAV18077.  
 XX  
 PS Claim 9; Page 22-23; 33pp; English.  
 XX  
 CC This polypeptide comprises human RP1, a novel member of the EBI  
 CC family which is associated with T cell activation, and which binds  
 CC to normal, but not carboxy end-truncated, adenomatous polyposis coli  
 CC (APC) protein. RP1 was identified using a differential mRNA  
 CC technique designed to isolate molecules which were differentially  
 CC expressed following combined stimulation by CD3 and CD28 trigger-  
 CC molecules. The full-length cDNA sequence was obtained by RACE PCR.  
 CC RP2 and partial RP3 clones (see AAV18078 and AAV18079) were also  
 CC identified. A claimed method for determining expression of an  
 CC aberrant form of APC protein involves contacting the sample with at  
 CC least part of an RP protein (especially RP1, RP2 or RP3) able to  
 CC bind to normal, but not aberrant APC protein, and measuring any  
 CC binding of the APC protein to the RP protein. Detecting aberrant  
 CC APC protein can be used for diagnosis of, or determining  
 CC Crohn's disease. A claimed method for screening for a disorder  
 CC characterised by inappropriate T cell activation involves contacting  
 CC a T cell sample with a nucleic acid encoding RP1, RP2 or RP3 and  
 CC determining any hybridisation. Such activation may indicate T-cell  
 CC lymphoma and this could be treated with RP inhibitors.  
 CC  
 XX  
 SQ Sequence 327 AA;

Query Match 60.08; Score 895.5; DB 19; Length 327;  
 Best Local Similarity 58.28; Pred. No. 3e-84;  
 Matches 171; Conservative 46; Mismatches 54; Indels 23; Gaps 5;

QY 1 MAVNVYSTVSTSENLSRDLAMVNDLSLNTYKTEQCSGAAYCGFMDLFGCVHLRK 60  
 DB 44 MAVNVYSTVSTSENLSRDLAMVNDLSLNTYKTEQCSGAAYCGFMDLFGCVHLRK 103  
 QY 61 VKFOAKLEHEYIHNKVLQAAFKKMGVDKIIPEVKLVKGFQDNFEFLQWFKKFFDANYD 120  
 DB 104 VKFOAKLEHEYIHNKVLQAAFKKMGVDKIIPEVKLVKGFQDNFEFLQWFKKFFDANYD 163  
 QY 121 GKDVLPPLARQGVAPPNPGDQIFNKSRL- - - - -IGTAVPQRTSPFG- - - - -PKNMQT 171  
 DB 164 GKDVLPPLARQGVAPPNPGDQIFNKSRL- - - - -IGTAVPQRTSPFG- - - - -PKNMQT 223

QY 172 SGRLSNVAAPCILRKNPSPARNGHETDAQIQLNOQLVDLKTVDGLERKDFYFSKLR 231  
 DB 224 AKRASSSG- - - - -SASKSDKDLQVYQLNQLVSLALAEVKEKEDFTFGKLR 273  
 QY 232 DIELICQHEHSENSPVISGIIILYATEGCFAPPEDELEE- -HOEE- -DODEY 281  
 DB 274 EIELICQHEHSENSPVISGIIILYATEGCFAPPEDELEE- -HOEE- -DODEY 327  
 RESULT 7  
 AAB43104  
 ID AAB43104 standard; Protein; 327 AA.  
 XX  
 AC AAB43104;  
 XX  
 DT 08-FEB-2001 (first entry)  
 XX  
 DE Human ORFX ORF2868 polypeptide sequence SEQ ID NO:5736.  
 XX  
 KW Human: open reading frame; ORFX; detection; cytosolic; hepatotropic;  
 KW vulnery; antipsoptic; antiparkinsonian; neurotropic; neuroprotective;  
 KW anticonvulsant; osteopathic; antiallergic; immunosuppressant; cardiac;  
 KW immunostimulant; thrombolytic; coagulant; vasotropic; antidiabetic;  
 KW hypotensive; dermatological; immunosuppressive; antineoplastic;  
 KW antiviral; antibacterial; antifungal; antineuritic; antithyroid;  
 KW antianemic; gene therapy; cancer; proliferative disorder; hypertension;  
 KW neurodegenerative disorder; osteoarthritis; graft vs host disease;  
 KW cardiovascular disease; diabetes mellitus; hypothyroidism; SCID; AIDS;  
 KW cholesterol ester storage; systemic lupus erythematosus; infection;  
 KW severe combined immunodeficiency; malaria; autoimmune disorder; asthma;  
 KW allergy; aplastic anaemia; nocturnal haemoglobinuria; burn; wound;  
 KW bone damage; cartilage damage; antineoplastic disease; coagulation;  
 KW thrombosis; contraceptive.  
 KW  
 XX  
 OS Homo sapiens.  
 XX  
 PN WO200058473-A2.  
 XX  
 PD 05-OCT-2000.  
 XX  
 PF 31-MAR-2000; 2000WO-US08621.  
 XX  
 PR 31-MAR-1999; 99US-0127607.  
 XX  
 PR 02-APR-1999; 99US-0127636.  
 XX  
 PR 05-APR-1999; 99US-0127728.  
 XX  
 PR 30-MAR-2000; 2000US-0540763.  
 XX  
 PA (CURA-) CURAGEN CORP.  
 XX  
 PI Shinkets RA, Leach M;  
 XX  
 DR WPI: 2000-602362/57.  
 XX  
 DR N-PSDB: AAC77313.  
 XX  
 PT Novel nucleic acids and peptides derived from open reading frame X,  
 PT useful for treating e.g. cancers, proliferative disorders,  
 PT neurodegenerative disorders and cardiovascular disease -  
 XX  
 PS Claim 11; Page 4899-4900; 5507pp; English.  
 XX  
 CC AAC74446 to AAC77606 encode the proteins given in AAB40237 to AAB43397,  
 CC which represent the human ORFX open reading frames 1 to 3161. The ORFX  
 CC sequences have activities such as: cytosolic; hepatotropic; vulnery;  
 CC antipsoptic; antiparkinsonian; neurotropic; neuroprotective;  
 CC osteopathic; anticonvulsant; antiallergic; immunosuppressant;  
 CC immunostimulant; cardiac; thrombolytic; coagulant; vasotropic;  
 CC antidiabetic; hypotensive; dermatological; immunosuppressive;  
 CC antineoplastic; antibacterial; antiviral; antifungal; antirheumatic;  
 CC antithyroid; and antianemic. The sequences can be used for determining  
 CC the presence of or predisposition to, or preventing or treating  
 CC pathological conditions associated with an ORFX-associated disorder. The  
 CC nucleic acids can be used to express ORFX proteins in gene therapy

CC vectors. The proteins and nucleic acids may be used to treat cancers,  
 CC proliferative disorders, neurodegenerative disorders, osteoarthritis,  
 CC graft vs host disease, cardiovascular disease, diabetes mellitus,  
 CC hypertension, hypothyroidism, cholesterol ester storage, systemic lupus  
 CC erythematosus, severe combined immunodeficiency (SCID), AIDS, viral,  
 CC bacterial or fungal infection, malaria, autoimmune disorders, asthma,  
 CC allergies, aplastic anaemia, burns, wounds, bone and cartilage damage,  
 CC nocturnal haemoglobinuria, anti-inflammatory disease; to enhance  
 CC coagulation; to inhibit thrombosis; and as a contraceptive.

CC Sequence 327 AA;

Query Match 60.0%; Score 895.5; DB 21; Length 327;  
 Best Local Similarity 58.2%; Pred. No. 3684;  
 Matches 171; Conservative 46; Mismatches 54; Indels 23; Gaps 5;

QY 1 MAVNVSTVSTSENLSRHMFLAVNDSLNLNTKIKDQSCGAAYCOFMDLFPGCYHLRK 60  
 DB 44 MAVNVSTSTQTSRHRDILAVNNDIVSLNTKVKVQLDSCGAAYCOFMDLFPGCISLKK 103  
 QY 61 VKFOAKLEHEYIHNFKVLQAAFKMGVDKIIPVEKIVKGFQDNFPIOMFKKFFPANDY 120  
 DB 104 VKFOAKLEHEYIHNFKVLQAAFKMGVDKIIPVEKIVKGFQDNFPIOMFKKFFPANDY 163  
 QY 121 GADYNPLAROGODVAPPPNPGDQIFNKSCLI--GTAVPQRTSPGK--PKNMOT 171  
 DB 164 GREYDVEAROGODAIIPDPDGEQIFNLPRKSHANSPTAGAAKSSPAKPGSTPSPSS 223  
 QY 172 SGRLSNVAPRCILRKPPSARNGCHETDAQILELNOQLVDLKTIVGLEKREFYFSKLK 231  
 DB 224 AKRASSG-----SKSKSDKLETVQVQLNDQVNSKLALGVEERDFYFGKLK 273  
 QY 232 DIEELQEHSESPVYSIGILVATEEGFAPPEDEIE--HQDE--DODEY 281  
 DB 274 EIEELQEHSESPVYSIGILVATEEGFAPPEDEIE--HQDE--DODEY 327

# RESULT 8

AAU30532 standard; Protein; 305 AA.

AC AAU30532;  
 DT 18-DEC-2001 (first entry)

DE Novel human secreted protein #1023.

XX Human; vaccination: gene therapy; nutritional supplement;  
 KM stem cell proliferation; haematopoiesis; nerve tissue regeneration;  
 KM immune suppression; immune stimulation; anti-inflammatory; leukaemia.

OS Homo sapiens.

PN WO200179449-A2.

PD 25-OCT-2001.

PF 16-APR-2001; 2001WO-US08656.

PR 18-APR-2000; 2000US-0552929.

PR 26-JAN-2001; 2001US-0770160.

PA (HYSE-) HYSEQ INC.

PI Tang YT, Liu C, Drmanac RT;

XX WPI; 2001-611725/70.

PT Nucleic acids encoding a range of human polypeptides, useful in genetic  
 PT vaccination, testing and therapy -

PS Claim 20; Page 306; 765pp; English.

CC The invention relates to novel human secreted polypeptides. The  
 CC polypeptides and antibodies to the polypeptides are useful for  
 CC determining the presence of or predisposition to a disease associated  
 CC with altered levels of polypeptide. The polypeptides are also useful for  
 CC identifying agents (agonists and antagonists) that bind to them. Cells  
 CC expressing the proteins are useful for identifying a therapeutic agent  
 CC for use in treatment of a pathology related to aberrant expression or  
 CC physiological interactions of the polypeptide. Vectors comprising  
 CC the nucleic acids encoding the polypeptides and cells genetically  
 CC engineered to express them are also useful for producing the proteins.  
 CC The proteins are useful in genetic vaccination, testing and  
 CC therapy, and can be used as nutritional supplements. They may be used to  
 CC increase stem cell proliferation; to regulate haematopoiesis; and in  
 CC bone, cartilage, tendon and/or nerve tissue growth or regeneration;  
 CC immune suppression and/or stimulation; as anti-inflammatory agents; and  
 CC in treatment of leukaemias. AAU29510-AAU33304 represent the amino acid  
 CC sequences of novel human secreted proteins of the invention.

CC Sequence 305 AA;

Query Match 57.4%; Score 857; DB 22; Length 305;  
 Best Local Similarity 61.8%; Pred. No. 2,7e-80;  
 Matches 175; Conservative 35; Mismatches 59; Indels 14; Gaps 6;

QY 1 MAVNVSTVSTSENLSRHMFLAVNDSLNLNTKIKDQSCGAAYCOFMDLFPGCYHLRK 60  
 DB 35 MAVNVSTVSTSENLSRHMFLAVNDSLNLNTKIKDQSCGAAYCOFMDLFPGCYHLRK 94  
 QY 61 VKFOAKLEHEYIHNFKVLQAAFKMGVDKIIPVEKIVKGFQDNFPIOMFKKFFPANDY 120  
 DB 95 VKFOAKLEHEYIHNFKVLQAAFKMGVDKIIPVEKIVKGFQDNFPIOMFKKFFPANDY 154  
 QY 121 GADYNPLAROGODVAPPPNPGDQIFNKSCLI--GTAVPQRTSPGKPMOTSGRLSNV 178  
 DB 155 GADYNPLAROGODVAPPPNPGDQIFNKSCLI--GTAVPQRTSPGKPMOTSGRLSNV 209  
 QY 179 APPCILRKPPSARNGCHETDAQILELNOQLVDLKTIVGLEKREFYFSKLDELICQ 238  
 DB 210 GRLGVVKKN-PGVNGDDDE-AELMOOGORIKNL-LFEDLGGERGFYFGKLRLNIELICQ 266  
 QY 239 EHESENSPVYSIGILVATEEGFAPPEDEIEHQDEDDDEY 281  
 DB 267 ENGENDPVQLRIVDILYATDEGFIID----EGGQDEDEY 305

# RESULT 9

ABB59772 standard; Protein; 291 AA.

AC ABB59772;

DT 26-MAR-2002 (first entry)

DE Drosophila melanogaster polypeptide SEQ ID NO 6108.

XX Drosophila; developmental biology; cell signalling; insecticide;  
 KM pharmaceutical.

OS Drosophila melanogaster.

PN WO200171042-A2.

PD 27-SEP-2001.

PF 23-MAR-2001; 2001WO-US09231.

PR 23-MAR-2000; 2000US-191637P.

PR 11-JUL-2000; 2000US-0614150.

PA (PEKE ) PE CORP NY.

PI Venter JC, Adams M, Li PMD, Myers EW;

DR WPI: 2001-656860/75.  
DR N-PSDB: ABL03875.  
PT New isolated nucleic acid detection reagent for detecting 1000 or more  
PT genes from Drosophila and for elucidating cell signalling and cell-cell  
PT interactions -  
PS Disclosure: SEQ ID NO 6108; 21pp + Sequence Listing; English.  
XX  
CC The invention relates to an isolated nucleic acid detection reagent  
CC capable of detecting 1000 or more genes from Drosophila. The invention is  
CC useful in developmental biology and in elucidating cell signalling and  
CC cell-cell interactions in higher eukaryotes for the development of  
CC insecticides, therapeutics and pharmaceutical drugs. The invention  
CC discloses genomic DNA sequences and the encoded proteins  
CC sequences (AB101840-AB16175) and the encoded proteins  
CC (AB57737-AB72072).  
CC The sequence data for this patent did not form part of the printed  
CC specification, but was obtained in electronic format directly from WIPO  
CC at ftp.wipo.int/pub/published\_pct\_sequences.  
XX  
SQ Sequence 291 AA:  
Query Match 52.4%; Score 782; DB 22; Length 291;  
Best Local Similarity 53.2%; Pred. No. 1.6e-72;  
Matches 160; Conservative 41; Mismatches 62; Indels 38; Gaps 7;  
OY 1 MAVNVYTSVTSSENLRHDMALWVNDLSHLNNTKTEQLCSGAAQCFMDLPPGCVHLRK 60  
DB 1 MAVNVYTSVTSSENLRHDMALWVNDLSHLNNTKTEQLCSGAAQCFMDLPPGCVHLRK 60  
OY 61 VKFOAKLEHEHYIHNFKVLOAFKMGVDKIIPEVKLVKGFQDNFEIOWFKKFPDAND 120  
DB 61 VKFRINLEHYIQNKILQAGFKKMSVDKIIPIDKLVKGRFDNFEIOWFKKFPDAND 120  
OY 121 GKDYNPLAROGQDVAPPNPGDQIFNKSKLIGTAV-----PORTSPT 164  
DB 121 GRDYDASAVREG---APMGFGS---GAVKSLPGTAASGVSSSYRRGSPATTPAMTSAY 173  
OY 165 GPKNMQTSGRLSNVAP-----PC-----ILKRNPSARNGGHETDQILLENQQLVDKL 214  
DB 174 KPTYSKVLPRTNNAPASRINACANSTGTVKKNVDS---NSVNNQQLIEMSNQVMDMRI 229  
OY 215 TVDGLKEKRDYFSKLRDIELCOE--HESENSPVISGIILYATEEGFAPPEDEIEEH 273  
DB 230 NLEGLEKERDYEFSKLRDIELCOEADDEAHPITIOKILIDILYATEDEGFAPPDDAPPEDE 289  
OY 274 Q 274  
DB 290 E 290  
RESULT 10  
ABB6121  
ID ABB6121 standard; Protein: 291 AA.  
AC ABB6121;  
XX  
DT 26-MAR-2002 (first entry)  
XX  
DE Drosophila melanogaster polypeptide SEQ ID NO 25155.  
XX  
KW Drosophila: developmental biology; cell signalling; insecticide;  
KW pharmaceutical.  
XX  
OS Drosophila melanogaster.  
XX  
PN MO200171042-A2.  
XX  
PD 27-SEP-2001.  
XX  
PF 23-MAR-2001; 2001WO-US09231.  
XX

PR 23-MAR-2000; 2000US-191637P.  
PR 11-JUL-2000; 2000US-0614150.  
XX  
XX (PEKE ) PE CORP NY.  
PA Venter JC, Adams M, Li PWD, Myers EW;  
PI  
DR WPI: 2001-656860/75.  
DR N-PSDB: ABL10224.  
XX  
CC New isolated nucleic acid detection reagent for detecting 1000 or more  
CC genes from Drosophila and for elucidating cell signalling and cell-cell  
CC interactions -  
PS Disclosure: SEQ ID NO 25155; 21pp + Sequence Listing; English.  
XX  
CC The invention relates to an isolated nucleic acid detection reagent  
CC capable of detecting 1000 or more genes from Drosophila. The invention is  
CC useful in developmental biology and in elucidating cell signalling and  
CC cell-cell interactions in higher eukaryotes for the development of  
CC insecticides, therapeutics and pharmaceutical drugs. The invention  
CC discloses genomic DNA sequences (AB101840-AB16175) and the encoded proteins  
CC sequences (AB101840-AB16175) and the encoded proteins  
CC (AB57737-AB72072).  
CC The sequence data for this patent did not form part of the printed  
CC specification, but was obtained in electronic format directly from WIPO  
CC at ftp.wipo.int/pub/published\_pct\_sequences.  
XX  
SQ Sequence 291 AA:  
Query Match 52.4%; Score 782; DB 22; Length 291;  
Best Local Similarity 53.2%; Pred. No. 1.6e-72;  
Matches 160; Conservative 41; Mismatches 62; Indels 38; Gaps 7;  
OY 1 MAVNVYTSVTSSENLRHDMALWVNDLSHLNNTKTEQLCSGAAQCFMDLPPGCVHLRK 60  
DB 1 MAVNVYTSVTSSENLRHDMALWVNDLSHLNNTKTEQLCSGAAQCFMDLPPGCVHLRK 60  
OY 61 VKFOAKLEHEHYIHNFKVLOAFKMGVDKIIPEVKLVKGFQDNFEIOWFKKFPDAND 120  
DB 61 VKFRINLEHYIQNKILQAGFKKMSVDKIIPIDKLVKGRFDNFEIOWFKKFPDAND 120  
OY 121 GKDYNPLAROGQDVAPPNPGDQIFNKSKLIGTAV-----PORTSPT 164  
DB 121 GRDYDASAVREG---APMGFGS---GAVKSLPGTAASGVSSSYRRGSPATTPAMTSAY 173  
OY 165 GPKNMQTSGRLSNVAP-----PC-----ILKRNPSARNGGHETDQILLENQQLVDKL 214  
DB 174 KPTYSKVLPRTNNAPASRINACANSTGTVKKNVDS---NSVNNQQLIEMSNQVMDMRI 229  
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DB 230 NLEGLEKERDYEFSKLRDIELCOEADDEAHPITIOKILIDILYATEDEGFAPPDDAPPEDE 289  
OY 274 Q 274  
DB 290 E 290  
RESULT 11  
AAG13505  
ID AAG13505 standard; Protein: 286 AA.  
AC AAG13505;  
XX  
DT 17-OCT-2000 (first entry)  
XX  
DE Arabidopsis thaliana protein fragment SEQ ID NO: 13024.  
XX  
KW Protein identification; signal transduction pathway; metabolic pathway;  
KW hybridisation assay; genetic mapping; gene expression control; promoter;  
KW termination sequence.  
XX

OS Arebidopsis thaliana.  
 XX  
 PN EPI033405-A2.  
 XX  
 PD 06-SEP-2000.  
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 PF 25-FEB-2000; 2000EP-0301439.  
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 PR 25-FEB-1999; 990S-0121825.  
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PR 28-OCT-1999; 99US-0161920.  
PR 28-OCT-1999; 99US-0161992.  
PR 28-OCT-1999; 99US-0161993.  
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Query Match 33.2%; Score 496; DB 21; Length 286;  
Best Local Similarity 36.3%; Pred. No. 7.7e-43;  
Matches 106; Conservative 58; Mismatches 80; Indels 48; Gaps 6;

QY 15 LSRHDM LAMVND SLH LNTYKT IEOLCS GAAYCO FMDL FPGCVHLRKVFQAKLEHEYIHN 74  
DB 7 VGRNEIILSMINDRLHLNLSRIEEAASGA VQCM LDMTFPGVVP MHKVNF EAKNEVEMION 66  
QY 75 FKVLQAAKKKGVKDIIV EKVGVKGFODNFEFIOMFKKFFDANYDG---KDYNFL--LA 129  
DB 67 YKVMQEVETFKLITKRP LRVKRPDNL EFLQWLEKRFCD SINGJMNENYNPERRS 126  
QY 130 RQGDVAPPPRGDQIFNKS---KKLIGTAVPQRTSPGPKNMOTSGRLSNVAPPCILRK 186  
DB 127 RGRREKSY---KGS SKIKRSIQTNMHPVATSNKPPAGPKOAKSHG----- 170  
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DB 171 -----IGGSSNSA VQALSKVEVDLKVSV DLEKERDFYFSKLNDIELICOTPELDDLP 225  
QY 247 VISGIGLILYATE-----EGRAPPEDEIEHQOEDD 279  
DB 226 IIVAVKILYATDANESVLEAEQCLINOSLGLEGE EEBGEBEEEEEEEEEE 277

RESULT 12  
ID AAG13504 standard; Protein; 287 AA.  
XX AAG13504;  
AC AAG13504;

XX 17-OCT-2000 (first entry)  
XX Arabidopsis thaliana protein fragment SEQ ID NO: 13023.  
DE Protein identification; signal transduction pathway; metabolic pathway;  
XX hybridisation assay; genetic mapping; gene expression control; promoter;  
KW termination sequence.  
XX Arabidopsis thaliana.  
OS EP1033405-A2.  
XX  
XX PD 06-SEP-2000.  
XX  
XX 25-FEB-2000; 2000EP-0301439.  
XX  
XX 25-FEB-1999; 99US-0121825.  
PR 05-MAR-1999; 99US-0123180.  
PR 09-MAR-1999; 99US-0123548.  
PR 23-MAR-1999; 99US-0125788.  
PR 25-MAR-1999; 99US-0126264.  
PR 29-MAR-1999; 99US-0126785.  
PR 01-APR-1999; 99US-0127462.  
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PR 18-JUN-1999; 99US-0139461.  
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PR 18-JUN-1999; 99US-0139463.





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PR 16-SEP-1999; 990S-0154039.  
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PR 18-OCT-1999; 990S-0159584.  
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PR 21-OCT-1999; 990S-0160767.  
PR 21-OCT-1999; 990S-0160768.  
PR 21-OCT-1999; 990S-0160770.  
PR 21-OCT-1999; 990S-0160814.  
PR 21-OCT-1999; 990S-0160815.  
PR 22-OCT-1999; 990S-0160980.  
PR 22-OCT-1999; 990S-0160981.  
PR 22-OCT-1999; 990S-0160989.  
PR 25-OCT-1999; 990S-0161404.  
PR 25-OCT-1999; 990S-0161405.  
PR 25-OCT-1999; 990S-0161406.  
PR 26-OCT-1999; 990S-0161359.  
PR 26-OCT-1999; 990S-0161360.  
PR 26-OCT-1999; 990S-0161361.  
PR 28-OCT-1999; 990S-0161920.  
PR 28-OCT-1999; 990S-0161922.  
PR 28-OCT-1999; 990S-0161993.  
PR 29-OCT-1999; 990S-0162142.

Query Match 33.2%; Score 496; DB 21; Length 293;  
Best Local Similarity 36.3%; Pred. No. 8643;  
Matches 106; Conservative 58; Mismatches 80; Indels 48; Gaps 6;

OY 15 LSRHMDLAVNVDLSNLTNTKRIEDLCSGAYCQFMHLPFGCVHLKRVKQAKLHEHYIIN 74  
DB 14 VGRNELLSMINRHLNLTLEIAASGAVQCOMLDMTFPGVPMHAKNEKYEYEMION 73  
OY 75 FYVLOAFKKMGVDKLIPEKLVKGFQDNFEFIOWFKKFPFANYG--KDYNPL--LA 129  
DB 74 YKVMQEVFTKLTITKPLEVNRLLKGRPLDNLFEFLQWLKRFCDISINGCINMENTINPVERRS 133

OY 130 RQGQVAPPPNPQDITFNKS---KKLIGTAVPQRTSPGPKMKQTSGRLSNVAPPCILRK 186  
DB 134 RQGREKSV---KGSSKIKSLQTNMHHPPVATSNKPPAGPKQAKSHG----- 177  
OY 187 NPPSARNGGHEHDAQLELNLQOLVDLKLTPDLEKERDFFFSKLDIELICQHESENSE 246  
DB 178 -----IGGSSNSAEVQALSKVEDLKVSVDLEKERDFFFSKLDIELICQTPPELDLP 232  
OY 247 VISGIIIGLYATE-----EGFAPPEDEIEHQEDQD 279  
DB 233 IVVAVKILYATDANESVLEAQECLNLSLGEYEECKEHEEEHEEEEEE 284  
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AAG47797  
ID AAG47797 standard; Protein; 316 AA.  
XX AAG47797;  
AC  
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DT 18-OCT-2000 (first entry)  
XX  
DE Arabidopsis thaliana protein fragment SEQ ID NO: 60282.  
XX  
KW Protein identification; signal transduction pathway; metabolic pathway;  
KW hybridisation assay; genetic mapping; gene expression control; promoter;  
KW termination sequence.  
XX Arabidopsis thaliana.  
XX  
PN EPI033405-A2.  
XX  
PD 06-SEP-2000.  
XX  
XX  
PF 25-FEB-2000; 2000EP-0301439.  
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PR 25-FEB-1999; 990S-0121825.  
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PR 25-MAR-1999; 990S-0126264.  
PR 29-MAR-1999; 990S-0126785.  
PR 01-APR-1999; 990S-0127462.  
PR 06-APR-1999; 990S-0128234.  
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PR 30-APR-1999; 990S-0132048.  
PR 04-MAY-1999; 990S-0133407.  
PR 04-MAY-1999; 990S-0133484.  
PR 05-MAY-1999; 990S-0133485.  
PR 06-MAY-1999; 990S-0132486.  
PR 06-MAY-1999; 990S-0132487.  
PR 07-MAY-1999; 990S-0132863.  
PR 11-MAY-1999; 990S-0134256.  
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PR 14-MAY-1999; 990S-0134219.  
PR 14-MAY-1999; 990S-0134221.  
PR 14-MAY-1999; 990S-0134370.  
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PR 21-MAY-1999; 990S-0135353.  
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PR 25-MAY-1999; 990S-0136021.  
PR 27-MAY-1999; 990S-0136392.  
PR 28-MAY-1999; 990S-0136782.  
PR 01-JUN-1999; 990S-0137222.  
PR 03-JUN-1999; 990S-0137528.  
PR 04-JUN-1999; 990S-0137502.

PR 07-JUN-1999; 99US-0137724.  
PR 08-JUN-1999; 99US-0138094.  
PR 10-JUN-1999; 99US-0138540.  
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PR 18-JUN-1999; 99US-0139461.  
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PR 22-JUN-1999; 99US-0139899.  
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PR 29-JUN-1999; 99US-0140991.  
PR 30-JUN-1999; 99US-0141287.  
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PR 01-JUL-1999; 99US-0142154.  
PR 02-JUL-1999; 99US-0142055.  
PR 06-JUL-1999; 99US-0142390.  
PR 08-JUL-1999; 99US-0142803.  
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PR 03-AUG-1999; 99US-0147038.  
PR 04-AUG-1999; 99US-0147204.  
PR 04-AUG-1999; 99US-0147302.  
PR 05-AUG-1999; 99US-0147192.  
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PR 06-AUG-1999; 99US-0147303.

PR 06-AUG-1999; 99US-0147416.  
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PR 11-AUG-1999; 99US-0148319.  
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PR 28-OCT-1999; 99US-0161993.  
PR 29-OCT-1999; 99US-0162142.

Query Match 32.4%; Score 483; DB 21; Length 316;  
Best local Similarity 35.3%; Pred. No. 2e-41;  
Matches 108; Conservative 63; Mismatches 89; Indels 46; Gaps 7;





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